



THE UNITED STATES PATENT AND TRADEMARK OFFICE

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APPLICANT: NORMAN K. SPROCH

DOCKET NO.: 0268P0342

SERIAL NO.: 09/287,307

EXAMINER: THAI PHAN

FILED: 04/07/1999

ART UNIT: 2123

TITLE: METHOD FOR THE CHARACTERIZATION OF THE THREE-DIMENSIONAL
STRUCTURE OF PROTEINS EMPLOYING MASS SPECTROMETRIC
ANALYSIS AND COMPUTATIONAL FEEDBACK MODELING

RECEIVED

APPELLANT'S BRIEF ON APPEAL

AUG 20 2003

Technology Center 2100

(1) Real Party in Interest

The inventor, Norman K. Sproch, is the real party in interest in this case.

(2) Related Appeals and Interferences

No other appeals or interferences are known to appellant or to the appellant's legal representative that would have any bearing on the Board's decision in this appeal.

(3) Status of Claims

The present application is a Continuation-In-Part of U.S. Patent Application Ser. No. 08/569,358, filed on December 08, 1995, now abandoned.

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Claims 1-18 are pending. All claims are rejected under 35 U.S.C. Sec. 102(e) as anticipated by Dunkel, U.S. Patent No. 5,572,125. The rejection of all pending claims is being appealed.

(4) Status of Amendments

No amendment was submitted after final rejection.

(5) Summary of Invention

The invention is a method for characterizing the three-dimensional structure of a large molecule (e.g., a protein molecule) comprising (1) mixing a small molecule with a large molecule (large and small molecules are defined on page 18, lines 9-19) so that the small molecule binds non-covalently to the large molecule to form a large molecule-small molecule complex (pages 35-37; Figs. 8-11), (2) performing electrospray ionization mass spectrometry (ES-MS) to obtain the spectrum of the large molecule-small molecule complex (page 19, line 21 through page 34; Figs. 1-7; for additional background on ES-MS, see U.S. Patent 5,504,327), (3) repeating the first two steps with additional different small molecules (pages 35-37; Figs. 17A-17C), and (4) using the spectrum so obtained to characterize the three-dimensional structure of the large molecule (pages 38-45).

In one preferred embodiment, ES-MS data is used to calculate the binding constant (K_B) for the binding of the small molecule to

the large molecule (pages 38-45; Fig. 21), the aforementioned mixing and ES-MS steps are repeated with additional different small molecules and the heat of formation (ΔH_f) for the binding of each of the small molecules to a selected residue on the surface of the large molecule is calculated (pages 41-43), the heat of formation (ΔH_f) for the binding of the small molecules to other selected residues on the surface of the large molecule is calculated (pages 41-45), the experimentally determined binding constant (K_B) is compared with the calculated heats of formation (Δh_f) (pages 43-45), and these comparisons are used to characterize the three-dimensional structure of the protein (pages 43-45). The three-dimensional molecular model elucidated through these comparisons can then further refined using experimental/computational feedback modeling (page 51, line 11 through page 55 ; Fig. 22).

(6) Issues

Whether Claims 1-18 were properly rejected under 35 U.S.C. Sec. 102(e) as anticipated by Dunkel, U.S. Patent No. 5,572,125.

(7) Grouping of Claims

The appellant believes that all claims should stand or fall together with respect to the prior-art rejection.

(8) Argument

The following argument is based in large part on the remarks presented to the Examiner in the appellant's response to the last Office Action.

Issue on Appeal- Rejection of Claim 1-18 Under 35 U.S.C. Sec. 102(e)

The appellant respectfully submits that there are at least two important differences between the invention claimed in the present application and that described in the U.S. Patent No. 5,572,125 (the Dunkel patent). Those differences include: (1) Dunkel's patent does not, indeed it cannot, describe a method for characterizing the three-dimensional structure of a large molecule as claimed by the appellant; and (2) While Dunkel's invention discusses the use of "complex molecules," nowhere does it disclose or suggest the appellant's claimed method of electrospray ionization mass spectrometry of molecule complexes.

As described throughout his patent, Dunkel's invention clearly is limited to a method for the correction of various types of signal distortion found in Fourier Transform Nuclear Magnetic Resonance Spectrometers. Indeed, his application of computer simulations using Monte Carlo computational methods is directed specifically toward the improvement of signal-to-noise ratios for the purpose of improving the spectra obtained.

In contrast, the present invention is based on two completely independent systems, one experimental and one computational, linked in such a way as to provide a 3-D, quantum mechanically correct, computer molecular model or image that, through computational methods, reveals the conformation of a protein (or other type of large molecule). Therefore, in the broadest sense, the appellant's invention represents the initial development of a system combining experimental and computational methods allowing for experimental data to be presented as a computer synthesized, chemically correct molecular image. In other words, the present invention describes a method for characterizing an experimental 3-D structure of a molecule being analyzed. Thus, the appellant's invention is completely unrelated to Dunkel's method for improving spectral data signal-to-noise ratios using Fourier transformation methods.

Turning to the specific reasons for rejection listed in the last Office Action, the Examiner begins by stating that "[Dunkel's] method includes the steps of mixing molecules to form a mixed solution of *complex molecules* for analysis" (emphasis added). In contrast, both independent claims (1 and 7) of the present application specifically recite "mixing a small molecule with a large molecule...to form a large molecule-small molecule complex." In other words, both claims 1 and 7 specifically recite molecule complexes, not "a mixed solution of *complex*

molecules." Since a "complex molecule" is not the same as a "molecule complex," Dunkel cannot anticipate the present invention as claimed.

With further regard to claim 1, the Examiner then states that Dunkel's method involves "performing electrospray ionization mass spectroscopy to obtain spectroscopic data of the molecule complexes." Despite the Examiner's general citation of several columns and sections, there is not a single mention of electrospray ionization mass spectrometry anywhere in Dunkel's patent. Thus, it is respectfully submitted that Dunkel does not describe the use of this type of spectrometry in his method. But even if it did, Dunkel's patent only deals with the correction of spectrum data, not the characterization of a 3-D structure based on the spectroscopic analysis of large and small molecule complexes as described and claimed by the appellant.

Lastly, the Examiner states that claim 1 is anticipated because Dunkel's method involves "repeating the procedure steps above if [sic] necessarily in order to obtain a good resolution for characterizing [the] 3-D structure [of] molecules." If understood correctly, this assertion by the Examiner is perplexing, as nowhere does Dunkel describe or suggest the characterization of the three-dimensional structure of large molecules by his method. In fact, the only "3-D" aspect of

Dunkel's method is displayed in Figs. 18a-f, 21a-f, and 24a-d, which show three-dimensional graphs of spectral data, not three dimensional structures of large molecules. These graphs of spectral data have absolutely no connection to a computationally synthesized, quantum mechanically correct, computer image of a molecule.

If anything, Dunkel's method is only useful for correctly predicting atom connectivity, not how the atoms are arranged in space. As one skilled in the chemical arts readily recognizes, a protein's (or other large molecule's) 3-D structure cannot be obtained solely from its atom connectivity.

Regarding independent claim 7, Dunkel only discloses a method and/or machine for acquiring, analyzing and correcting spectral and imaging data for the sole purpose of improving signal-to-noise values, thus yielding spectra with improved signal. Moreover, Dunkel's computer simulations are based entirely on Monte Carlo simulation with only unidirectional feedback (see col. 37, line 29 to col. 42, line 43).

Furthermore, as discussed above for claim 1, the Examiner's reliance on the disclosure in Dunkel of a "mixed solution of complex molecules" to anticipate the claims of the present invention is clearly based on a misreading of the "mixing a

small molecule with a large molecule...to form a large molecule-small molecule complex" limitation found in claim 7. Again, Dunkel's invention can provide no insight into, nor does it suggest any possible way of elucidating, the three-dimensional structure of large molecules (and especially not 3-D structures bases on spectrometry of large molecule/small molecule complexes).

Furthermore, Dunkel does not anticipate claim 7 based on, as stated by the Examiner, "bonding strength based on spectroscopic data in simulation model which would include bonding strength, bond energy, etc., as known for those skilled in the spectroscopy analysis." Bonding strength and bond energy cannot be obtained from the method described by Dunkel. This is because these thermodynamic properties must be calculated using suitable, specialized programs based on ab initio or semi-empirical quantum mechanical methods (all of which are outlined in the present application but are not even mentioned by Dunkel).

In fact, Dunkel's use of the stochastic Monte Carlo method would not allow the calculation of thermodynamic properties, even if Dunkel wished to use the Monte Carlo method for something other than noise reduction. In addition, Dunkel does not anticipate, as stated by the examiner, "data model being corrected to improve a selected residue on the molecule, and repeating the procedure

steps above if necessarily in order to obtain a good resolution for characterizing 3-D structure molecules." First, a residue is a specific chemical term used to describe one amino acid in a polypeptide chain or protein. Dunkel makes no mention of 'residues', and even if he had, his invention would only refer to the linear connectivity (not 3-D structure) of these residues in a chain of amino acids.

Dunkel also does not disclose "characterizing 3-D structure molecules." Dunkel is merely using an n-dimensional mathematical model, based on estimates, to simulate a signal from which the noise may be extracted. Thus, a 3-D mathematical model, as described by Dunkel, has nothing to do with the 3-D structure or 3-D computer image of a molecule.

In view of the clear differences between Dunkel and both independent claims of the present invention, the appellant respectfully submits the claim 1 and claim 7 are not in any way anticipated. Since each independent claim is clearly distinguishable from Dunkel, each dependent claim would also be so distinguished. Therefore, the appellant submits that all dependent claims would also be allowable as presently written.

However, in the interest of further clarifying the present invention, the following discussion addresses the Examiner's reasons for rejection of the dependent claims.

Regarding claims 2 and 8, Dunkel's invention uses a "computerized data processing system including plurality of means for performing steps, such as processing means for computing error data, phase shift data, etc., memory for storing computational results," solely for improving signal-to-noise ratios. This is accomplished by reducing noise through the specific application of Fourier transformation methods, resulting in an improved spectrum. As mentioned previously, this data correction method has nothing to do with the present invention, which produces an accurate three-dimensional representation of a molecule through the use of "feedback modeling."

In fact, the appellant's method of "feedback modeling" would allow for the elucidation of new data not obtainable from the spectra alone, by, for example, the application of quantum mechanically accurate computer modeling programs. Dunkel is not able to use his invention to provide new data not already contained in the Fourier transformation acquisition; he can only reduce the noise found in existing data. Moreover, while the modeling programs referred to in the present application have been well documented in the literature, they have not been used previously in a feedback loop as described. Thus, the present invention is clearly new.

With further regard to claim 2, the Examiner states that "Dunkel also anticipates simulating the model to predict error and correct the model using feedback loop as claimed." Again, Dunkel's use of Monte Carlo methods represent a stochastic approach for the purpose of eliminating noise to produce an improved spectrum. In fact, he is not "simulating the model;" he is simulating the experimental system, based on initial estimates. In other words, the feedback loop used by Dunkel is based on feedback from the simulation to the experimental data, which is, in effect, a unidirectional feedback. Dunkel does not allow for any manipulation or modification of the chemical system being analyzed prior to, during, or after the simulation.

In contrast, the present invention uses a bi-directional feedback, allowing for computer-controlled manipulation of experimental parameters, and allowing for chemical changes to be made in the chemical system during analysis. This results in new experimental data that is returned to the computational system for further computation, the results of which control changes in the experimental chemical system, which allows for the elucidation of new data (and not merely a reduction in noise) not obtainable by either system independently.

Regarding claims 3-6, the Examiner states that the "variety of complex molecules such as cholesterols, proteins and protein

complex structures" anticipates the appellant's claims. However, the Examiner again fails to recognize that molecule complexes, not "complex molecules," are claimed in the present invention. Thus, the fact that the Examiner considers cholesterol or a protein to be a "complex molecule" is irrelevant to present invention. Moreover, what exactly the Examiner means by "protein complex structures" is not understood, for nowhere does Dunkel disclose the ability to use his invention with proteins or complexes of proteins with small molecules. In fact, all of Dunkel's examples and data involve small bioorganic molecules, including cholesterol, Na_2MoO_4 , conotoxin, and 1,1-dimethyltetralin.

Moreover, Dunkel's only reference to proteins is with regard to protein sequencing (col. 33, lines 59-62), which cannot be construed to mean whole proteins as protein sequencing invariably involves the analysis of a series of small polypeptide fragments. Besides, protein sequencing provides only the linear connectivity of amino acids, not the 3-D conformation. Therefore, Dunkel's reference to proteins or "polymers" does not imply that he can apply his method to whole proteins, DNA, RNA or other large biomolecules. This is supported by the fact that (1) Dunkel uses a 500 MHz NMR spectrometer (col. 21, lines 30-40), which would not be sufficient for the analysis of a protein such as Cytochrome c (the molecule characterized by the appellant); and

(2) the largest molecule identified by Dunkel, conotoxin, is a small (22 amino acid) polypeptide, not a protein.

Regarding claims 9-15, again the Examiner fails to recognize that a molecular complex has an entirely different meaning from "complex molecules." Once more, Dunkel's invention is directed toward the reduction of noise to improve a data signal using Fourier transformation analysis and Monte Carlo simulations. Even if one could predict a 3-D structure with Dunkel's method as the Examiner suggests, only very small structures could be inferred, and then only through the application of basic chemical principles (none of which are described by Dunkel).

However, a likely 3-D structure cannot be inferred using basic chemical principles when large molecules or macromolecules, such as proteins, are being described. Instead, the 3-D structure of proteins must be determined using other experimental methods, none of which are mentioned by Dunkel. Thus, the present invention is clearly distinguishable from Dunkel because it describes and specifically claims a method for characterizing the 3-D structure of a large molecule.

Regarding claim 16, the examiner states that, "Dunkel anticipates bonding strength or binding energy of complex molecules such [that?] energy required to create a bond which would inherently

include heat of formation in the complex large molecules claimed." The appellant respectfully disagrees. Dunkel does not make any statements regarding bonding strength, binding energy, or heat of formation. Nowhere in his patent are these thermodynamic properties inferred nor calculated. Indeed, Dunkel only mentions "bond signals," i.e., spectral data, which have absolutely no connection to these thermodynamic properties.

Finally, regarding claims 17-18, the Examiner's interpretation of Dunkel's disclosure as describing a "plurality of complex molecules which would include and not limited to the claimed invention," is not correct. Again, a "plurality of complex molecules" has absolutely no connection to a molecular complex, which is key in understanding the present invention. Furthermore, Dunkel makes no disclosure regarding protein/small molecule complexes.

In summary, unlike the claims of the present application, there is no disclosure or suggestion in Dunkel of a method that elucidates the 3-D structure of macromolecules, proteins, DNA, or RNA based on spectrometry of molecular complexes resulting from the non-covalent interaction of a macromolecule or protein with a small molecule.

In view of the foregoing, the appellant respectfully submits that all pending claims recite allowable subject matter. Accordingly, the appellant believes that the Examiner erred in rejecting the claims and urges the Board to so hold.

Respectfully submitted,

A handwritten signature in black ink, appearing to read 'Gavin J. Milczarek-Desai', written in a cursive style.

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(9) Appendix

The claims involved in this appeal read as follows:

1. A method for characterizing the three-dimensional structure of a large molecule comprising the steps of:

(a) mixing a small molecule with a large molecule so that the small molecule binds non-covalently to the large molecule to form a large molecule-small molecule complex;

(b) performing electrospray ionization mass spectrometry to obtain the spectrum of the large molecule-small molecule complex;

(c) repeating steps (a)-(b) with additional different small molecules; and

(d) utilizing the spectra obtained in steps (a)-(c) to characterize the three-dimensional structure of the large molecule.

2. The large molecule characterization method of Claim 1, wherein the three-dimensional structure characterization of step (d) is carried out by feedback modeling according to the following steps:

(e) providing data processing means;

(f) providing data storage means;

(g) digitizing raw experimental data acquired according to steps (a)-(c);

(h) storing the digitized data in said data storage means;

(i) initializing and running a selected computer program on said data processing means for simulating the experiment performed in steps (a)-(c);

(j) comparing simulation data obtained from step (i) with the digitized data from the experiment performed in step (g);

(k) if the comparing step (j) produces a result outside a predetermined parameter, establishing a feedback loop and initiating an iterative subroutine whereby the computer simulation adjusts itself, in an incremental way, to fit the simulation to the experimental value, compares the result to the experiment after each computational step and feeds the experimental data back into the input loop of the computation until the result of the comparison of step (j) is within the predetermined parameter.

3. The large molecule characterization method of Claim 1, wherein the large molecule is selected from the group consisting of polypeptides, proteins, DNA, RNA, oligosaccharides, and polymers thereof.

4. The large molecule characterization method of Claim 1, wherein the small molecules include crown ethers, macrocyclic polyethers, cryptands, and/or polymers of these compounds, or any other suitable ligand or macrocyclic ligand.

5. The large molecule characterization method of Claim 2, wherein the large molecule is selected from the group consisting of polypeptides, proteins, DNA, RNA, oligosaccharides, and polymers thereof.

6. The large molecule characterization method of Claim 2, wherein the small molecules include crown ethers, macrocyclic polyethers, cryptands, and/or polymers of these compounds, or any other suitable ligand or macrocyclic ligand.

7. A method for characterizing the three-dimensional structure of a large molecule comprising the steps of:

(a) mixing a small molecule with a large molecule so that the small molecule binds non-covalently to the large molecule to form a large molecule-small molecule complex;

(b) performing electrospray ionization mass spectrometry to obtain the spectrum of the large molecule-small molecule complex;

(c) using the spectrum from step (b) to calculate the binding constant K_b for the binding of the small molecule complex;

(d) repeating steps (a)-(c) with additional different small molecules;

(e) calculating the heat of formation (ΔH_f) for the binding of each of the small molecules used in steps (a)-(d) to a selected residue on the large molecule;

(f) repeating step (e) for other selected residues on the large molecule;

(g) comparing the binding constants (K_b) calculated in steps (c) and (d) with the ΔH_f values calculated in steps (e) and (f); and

(h) utilizing the comparisons of step (g) to characterize the three-dimensional structure of the large molecule.

8. The large molecule characterization method of Claim 7, wherein said comparing step (g) is carried out by feedback modeling according to the following steps:

(i) providing data processing means;

(j) providing data storage means;

(k) digitizing raw experimental data acquired according to steps (a)-(d);

(l) storing the digitized data in said data storage means;

(m) initializing and running a selected computer program on said data processing means for simulating the three-dimensional structure of said large molecule according calculations performed in steps (e)-(f);

(n) comparing simulation data obtained from step (m) with the digitized data from the experiment performed in step (k);

(o) if the comparing step (n) produces a result outside a predetermined parameter, establishing a feedback loop and initiating an iterative subroutine whereby the computer

simulation adjusts itself, in an incremental way, to fit the simulation to the experimental value, compares the result to the experiment after each computational step and feeds the experimental data back into the input loop of the computation until the result of the comparison of step (n) is within the predetermined parameter.

9. The large molecule characterization method of Claim 7, wherein the comparisons of step (g) are utilized to identify the residue or residues on the surface of the protein molecule to which the small molecule is bound.

10. The large molecule characterization method of Claim 7, wherein the large molecule is selected from the group consisting of polypeptides, proteins, DNA, RNA, oligosaccharides, and polymers thereof.

11. The large molecule characterization method of Claim 7, wherein the small molecules include crown ethers, macrocyclic polyethers, cryptands, and/or polymers of these compounds, or any other suitable ligand or macrocyclic ligand.

12. The large molecule characterization method of Claim 8, wherein the large molecule is selected from the group consisting

of polypeptides, proteins, DNA, RNA, oligosaccharides, and polymers thereof.

13. The large molecule characterization method of Claim 8, wherein the small molecules include crown ethers, macrocyclic polyethers, cryptands, and/or polymers of these compounds, or any other suitable ligand or macrocyclic ligand.

14. The large molecule characterization method of Claim 9, wherein the large molecule is selected from the group consisting of polypeptides, proteins, DNA, RNA, oligosaccharides, and polymers thereof.

15. The large molecule characterization method of Claim 9, wherein the small molecules include crown ethers, macrocyclic polyethers, cryptands, and/or polymers of these compounds, or any other suitable ligand or macrocyclic ligand.

16. The large molecule characterization method of Claim 7, further comprising the step of using the heat of formation calculated in step (e) and calculating the heat of reaction (ΔH_{RXN}) for the binding of each of the small molecules used in steps (a)-(d) to a selected residue on the large molecule.

17. The large molecule characterization method of Claim 16, wherein the large molecule is selected from the group consisting of polypeptides, proteins, DNA, RNA, oligosaccharides, and polymers thereof.

18. The large molecule characterization method of Claim 16, wherein the small molecules include crown ethers, macrocyclic polyethers, cryptands, and/or polymers of these compounds, or any other suitable ligand or macrocyclic ligand.



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Alice B. Vanicek

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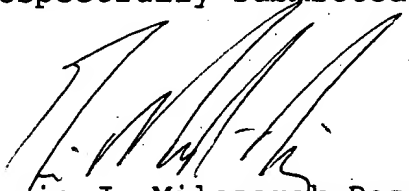
Dear Sir:

Pursuant to the provisions of 37 C.F.R. 1.192, the appellant is
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credit any overpayment associated with the filing of this Brief
on Appeal to our Deposit Account No. 04-1935.

Respectfully submitted,

A handwritten signature in black ink, appearing to read 'G. Milczarek-Desai', written over the typed name.

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